Blast Result

EXHIBITG



BLAST 2 SEQUENCES RESULTS VERSION BLASTN 2.2.2 [Dec-14-2001]

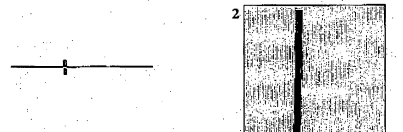
Match: 1 Mis match: 2 gap open: 5 gap extension: 2

x_dropoff: 50 expect: 10.000 wordsize: 11 Filter Align

Sequence gi Homo sapiens endothelial differentiation, sphingolipid G1 488: 194 protein-coupled receptor, 3 (EDG3), mRNA

Length 1137 (1...
1137)

Sequence | lcl|s: q 2



NOTE: The statistics (bitscore and expect value) is calculated based on the size of nr database

NOTE:If proteir translation is reversed, please repeat the search with reverse strand of the query sequence

Score = 44.9 pits (23), Expect = 0.17
Identities = 23/23 (100%)
Strand = Plus / Plus

Query: 423 caaaatgaggccttacgacgcca 445

Sbjetr 1 caaatgaggeettacgacgeca 23

CPU time: 0.05 user secs. 0.13 sys. secs 0.18 total secs.

Lambda K H
1.33 C.621 1.12

Gapped
Lambda K H
1.33 C.621 1.12

Matrix: blastn matrix:1 -2

Gap Penalties: Existence: 5, Extension: 2

Number of Hits to DB: 1

Blast Result

Number of Sequences: 0 Number of extensions: 1 Number of successful extensions: 1 Number of sequences better than 10.0: 1 length of query: 1137 length of database: 5,006,917,935 effective HSP length: 24 effective length of query: 1113 effective length of database: 4,991,889,975 effective sea ch space: 5555973542175 effective sea ch space used: 5555973542175 T: 0 A: 30 X1: 6 (11.5 b.ts) X2: 26 (50.0 bits) S1: 12 (23.8 pits) \$2: 20 (39.1 bits)